Tensor Networks and their use for Lattice Gauge Theories

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Tensor Network States are ansätze for the efficient description of quantum many-body systems. Their success for one dimensional problems, together with the fact that they do not suffer from the sign problem and can address the simulation of real time evolution, have turned them into one of the most promising techniques to study strongly correlated systems. In the realm of Lattice Gauge Theories they can offer an alternative to standard lattice Monte Carlo calculations, which are suited for static properties and regimes where no sign problem appears. The application of Tensor Networks to this kind of problems is a young but rapidly evolving research field. This paper reviews some of the recent progress in this area, and how, using one dimensional models as testbench, some fundamental milestones have been reached that may pave the way to more ambitious goals.

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1. Introduction

The term Tensor Network States (TNS) encompasses many different families of ansätze that can efficiently describe the state of a quantum many-body system. By construction, TNS families correspond to definite entanglement patterns. Quantum information theory provides tools to understand why they can be expected to be good ansätze for the physically relevant states, and some of the limitations connected to the simulation algorithms [1, 2, 3].

In the context of numerically investigating strongly correlated systems, TNS have demonstrated to be very powerful, specially for one dimensional problems, with the matrix product state (MPS) ansatz underlying the success of the density matrix renormalization group (DMRG) algorithm [4, 5, 6, 7], currently the most precise tool for the study of one-dimensional quantum many-body systems. The natural generalization of MPS to higher dimensions, Projected Entanglement Pair States (PEPS) [8] are good candidates to describe the physics of higher dimensional lattices. Another TNS ansatz, the MERA [9], can capture critical behavior and has recently been connected to a discrete realization of the AdS/CFT correspondence [10].

Lattice Gauge Theories (LGTs) offer challenging scenarios which can benefit from these techniques. The spatial dimensions and sizes of the systems amenable to TNS studies are still far from those solved by Monte Carlo simulations but, being free from the sign problem, tensor networks can be readily used for problems which more standard techniques cannot easily tackle. Some early works already combined the gauge invariance with TNS methods [11, 12, 13], and lately there has been a renewed interest on the topic, motivated in part by the prospects of quantum simulation, and also by the advances in the understanding and application of tensor networks.

During the last years, different teams have applied TNS algorithms to low dimensional lattice gauge theories from different perspectives, and it would be impossible to cover all these efforts in this limited space. The present discussion is thus restricted to studies in which the LGT is the regularization of a more fundamental quantum field theory in the continuum, and the ultimate goal is to solve the latter, much in the spirit of lattice QCD. Moreover, we focus on the use of TNS as ansatz for the physical states, in a Hamiltonian formulation of the theory.

Using one dimensional models as a testbed, different studies have demonstrated systematically that the MPS ansatz is well suited to describe the relevant physics in LGTs, and allows for very precise calculations of low energy and thermal properties, also in scenarios in which Monte Carlo techniques are hindered by the sign problem. This article reviews such results, which constitute necessary initial milestones before applying TNS to more ambitious LGT problems. The article is structured as follows. In section 2 the main TNS ideas and algorithms employed in these studies are discussed. Section 3 presents the models that have been used as testbench, and in section 4 we review the existing results. Finally, section 5 discusses the status and perspectives of the field.

2. Basic concepts

If we consider $N$ quantum systems, each one with finite physical dimension $d$, any quantum many body state can be exactly expressed in a tensor product basis by specifying $d^N$ coefficients, $|\Psi\rangle = \sum_{i_1,...,i_N=1}^{d} C_{i_1...i_N} |i_1...i_N\rangle$. We can think of the $C_{i_1...i_N}$ coefficients as the $d^N$ components of a $N$-legged tensor (which in this context simply denotes a multidimensional array). The exponen-
The growth of the dimension of the Hilbert space with the number of sites $N$ restricts the exact calculations to very small system sizes. However, we may consider restricted ansätze where the state coefficients can be expressed as the contraction of a certain network of smaller tensors, as illustrated in figure 1. Different structures of the network give rise to different TNS families, and naturally correspond to a different pattern of entanglement in the represented state.

While random states in the Hilbert space typically have close to maximum entanglement, ground states of physical local Hamiltonians behave in a very different manner. They often fulfill what is called the area law of entanglement. Namely the entropy of a subsystem grows only with the size of the boundary that separates it from the rest. In one spatial dimension, the area law has been rigorously proven to hold for ground states of local gapped Hamiltonians, and for critical cases it is known to be violated only logarithmically [14, 15].

This idea justifies the search for adequate families which, in terms of only a polynomial number of parameters and a structure based on the expected entanglement, are capable of capturing the essential features of the relevant states.

2.1 TNS in 1D: Matrix product states

The most popular tensor network, for which algorithms are the most efficient, and also the one that is best understood from a mathematical point of view, is the matrix product state family. A MPS is a state of the form $|\Psi\rangle = \sum_{i_1,\ldots,i_N} \text{tr}(A_{i_1}^1 A_{i_2}^2 \cdots A_{i_N}^N) |i_1 i_2 \ldots i_N\rangle$. Each $A_j^k$ is a matrix of dimensions $D \times D$ (equivalently $A_j^k$ is a rank-three tensor of dimensions $d \times D \times D$). In the case of open boundary conditions, the matrices for the first and last sites reduce to $D$-dimensional vectors, and the trace can be written as a product of matrices. The bond dimension $D$ determines the number of parameters in the ansatz, $d N D^2$. The entanglement between two halves of the chain is upper-bounded by $S(N/2)_{\text{max}} \propto \log D$. This shows explicitly that MPS satisfy a one-dimensional area law, which actually makes them expected good ansätze for ground states of local Hamiltonians. A single tensor can be used to parametrize a translationally invariant state, where $A_k = A$ for every site $k$, either for a finite periodic chain, or for an infinite system. With this ansatz, named uniform MPS (or uMPS), it is possible to directly work in the thermodynamic limit (see e.g. [2, 5] for reviews).

Global and local symmetries of the state can be encoded in properties of the individual tensors [16, 17, 18]. Investigating which classes of states can be represented by TNS with different symmetry restrictions is an active field of research. From the point of view of numerical calculations, explicitly imposing the gauge symmetry at the level of the individual tensors can reduce the number of variational parameters (and thus the computational cost), ensuring at the same time that the
ansatz is in the desired sector, for instance, the physical gauge-invariant subspace in the case of a
lattice gauge theory.

Different numerical methods exist to solve a given physical problem with MPS. The following
paragraphs briefly review the main aspects of the ones that have been so far applied to LGT prob-
lems. Many of these algorithms have been generalized to other TNS families, although in general
this involves much higher computational costs.

2.2 Variational optimization

In order to find a MPS approximation to the ground state of a local Hamiltonian, it is pos-
sible to variationally minimize the energy within the MPS family with fixed bond dimension by
sequentially optimizing the tensors in the ansatz one at a time, and iterating the procedure until
convergence. This variational strategy basically describes the extremely successful DMRG algo-

rithm and can easily be extended to obtain low lying excitations by searching for the lowest energy
state that is orthogonal to previously computed levels \[2, 5\].

In the case of uMPS in the thermodynamic limit, it is also possible to parametrize the exci-
tations as a single (or few) different tensors over the uniform background that corresponds to the
ground state. A state of well-defined momentum can be constructed as an appropriate superposition
of all possible translations of this construction, and the tensor parametrizing the excitation can also
be optimized variationally \[19, 20\].

2.3 Time evolution

It is also possible to find an approximation to the ground state using imaginary (or Euclidean)
time evolution. The most widely used strategy to evolve a MPS, in real or imaginary time, is based
on a Suzuki-Trotter expansion of the exponential evolution operator \[2, 5\]. The time is discretized
in small steps, whose corresponding evolution operators can be expressed as products of local terms
(e.g. two-body operators in the case of a nearest-neighbor model). The action of each such term on
the MPS yields in general a MPS of larger bond dimension, so the result needs to be truncated, a
step that may introduce a numerical error. This strategy is in practice limited to local Hamiltonians,
but more sophisticated algorithms exist that can also cope with long range interactions \[19\].

The algorithm is essentially the same for real or imaginary time evolution, but in the former
problem the entanglement in the physical state may grow very fast (up to linearly with the elapsed
time), which implies that the bond dimension required to keep an accurate description of the time-
dependent state as a MPS will increase exponentially. This in practice limits the feasibility of real
time simulations to short times or close to equilibrium situations.

2.4 Thermal states

Quantum many-body operators can be expressed in a tensor product basis in which every
element is the tensor product of one element of a local basis for each site of the system. If the
coefficients of the expansion have a MPS structure, we talk about a matrix product operator (MPO),
an extremely useful form of writing Hamiltonians and approximating evolution operators. Thermal
equilibrium and other mixed states are described by a density operator, and the MPO ansatz can be
applied to approximate them \[21, 22, 23\]. Moreover, it is known to do it efficiently in the case of
local Hamiltonians (also in higher dimensions) \[24\].
The positivity constraint, necessary to ensure that the MPO ansatz encodes a physical state, is not easy to impose at the level of the individual tensors. One way to overcome this problem is to use a more restricted MPO ansatz, referred to as purification, in which each of the tensors has a positive structure [21, 25]. This is equivalent to assuming a certain MPS form for a pure state that describes an enlarged system, and in the case of the thermal equilibrium states corresponds to a thermofield doubling. The purified state for inverse temperature $\beta$ is then, up to normalization, the result of applying the exponential operator $e^{-\beta H/2}$ to a maximally entangled state of system and ancilla. With the appropriate ordering of system and ancilla sites, such state is a trivial MPS, and the action of the exponential can be approximated using an algorithm for imaginary time evolution. This yields a very stable and widely used algorithm to find thermal states with TNS.

3. Models

One dimensional models are a natural testbench to investigate the performance of TNS numerical methods. The studies reported here have considered several models of fermions interacting with gauge fields. In the discrete formulation, fermions reside on the vertices of the lattice and gauge degrees of freedom on the links between them and the MPS ansatz can be formulated in a basis which is the tensor product of the local basis for fermionic and gauge degrees of freedom.

**Schwinger model.** One of the key example models is the Schwinger model, or QED in one spatial dimension. Arguably the simplest gauge theory with matter, it exhibits phenomena, as confinement and chiral symmetry breaking, common to more complex gauge theories, and a phase transition in presence of a background field (which is the equivalent in this case of a topological theta term). In the temporal gauge, and using Kogut-Susskind formulation with staggered fermions, the discretized Hamiltonian in the absence of background field reads [26],

$$H = -i \frac{a}{2} \sum_n \left( \phi_n^\dagger e^{i\theta_n} \phi_{n+1} - \text{h.c.} \right) + m \sum_n (-1)^n \phi_n^\dagger \phi_n + \frac{ag^2}{2} \sum_n L_n^2,$$

where $\phi_n^\dagger$ represents the creation operator of a spinless fermion on lattice site $n$, and $U_n = e^{i\theta_n}$ is the link operator between sites $n$ and $n+1$. $L_n$, canonical conjugate to $\theta_n$, corresponds to the electric field on the link. Gauss law appears as an additional constraint that has to be satisfied by physical states, namely $L_n - L_{n-1} = \phi_n^\dagger \phi_n - \frac{1}{2} [1 - (-1)^n]$. The parameters of the model are the fermion mass $m$ and the coupling $g$ (the only parameters in the continuum), and the lattice spacing $a$. It is convenient to rescale the Hamiltonian and work with adimensional parameters, $x = 1/(ag)^2$, $\mu = 2\sqrt{3m}/g$, such that the continuum limit corresponds to $x \to \infty$. The local Hilbert space basis for the fermionic sites can be labeled by the occupation of the mode, $n_m \in \{0, 1\}$ (for site $m$), while the basis elements for the links can be labeled by the integer eigenvalues of $L_n$, $\ell_m \in \mathbb{Z}$.

**Multiflavor Schwinger model.** A generalization of the model above consists in considering several different fermion flavors, each one with a different chemical potential $\kappa_f$. Except for certain specific choices of the parameters, conventional Monte Carlo techniques will encounter here a sign problem. The modified Hamiltonian can be written

$$H = -i \frac{a}{2} \sum_{n=0}^{N-2} \sum_{f=0}^{F-1} \left( \phi_{n,f}^\dagger e^{i\theta_n} \phi_{n+1,f} - \text{h.c.} \right) + \sum_{n=0}^{N-1} \sum_{f=0}^{F-1} (m_f (-1)^n + \kappa_f) \phi_{n,f}^\dagger \phi_{n,f} + \frac{ag^2N-2}{2} \sum_{n=0}^{N-1} L_n^2,$$

where $\phi_{n,f}^\dagger$ represents the creation operator of a spinless fermion with flavor $f$ on lattice site $n$. $U_n = e^{i\theta_n}$ is the link operator between sites $n$ and $n+1$. $L_n$, canonical conjugate to $\theta_n$, corresponds to the electric field on the link.
which can also be expressed in terms of adimensional parameters \( x = 1/(ag)^2 \), \( \mu_f = 2\sqrt{x}\kappa_f/g \). Gauss law now reads \( L_n - L_{n-1} = \sum_{f=0}^{F-1} \left[ \phi_n^+ \phi_{n,f} - \frac{1}{2} (1 - (-1)^n) \right] \), involving a sum over all fermion flavors. While the structure of the links does not change with respect to the previous case, the local Hilbert space for the fermionic site \( i \) is described by \( F \) occupation numbers corresponding to each flavor, \( \otimes_f |m_{n,f}\rangle \). The structure of the ground state in this case shows a series of transitions between phases with different imbalances \([27]\). Thus, it is an ideal problem to probe the performance of TNS in a simple scenario that yet is difficult for standard Monte Carlo methods.

**SU(2) in 1+1 D.** Another fundamental ingredient that a numerical technique has to work with is a non-Abelian gauge symmetry. The simplest model we can consider is a 1+1 dimensional SU(2) theory, for which the discrete Hamiltonian in the staggered fermion formulation reads

\[
H = \frac{1}{2a} \sum_{n=1}^{N-1} \sum_{\ell,\ell' = 1}^{2} \left( \phi_n^\dagger U_{n,\ell}^{\ell'} \phi_{n+1,\ell'} + \text{h.c.} \right) + m \sum_{n=1}^{N} \sum_{\ell=1}^{2} (-1)^n \phi_n^\dagger \phi_n^\tau + \frac{ag^2}{2} \sum_{n=1}^{N-1} J_n^2. \tag{3.3}
\]

Now the link operators \( U_{n,\ell}^{\ell'} \) are SU(2) matrices in the fundamental representation, and can be interpreted as rotation matrices. Gauss law is also a non-Abelian constraint, \( G_m^\tau |\Psi\rangle = 0, \forall m, \tau, \) with generators \( G_m^\tau = L_m - R_m - \sigma_m^\tau \), where \( Q_m^\tau = \sum_{\ell=1}^{2} \frac{1}{2} \phi_m^\dagger \phi_m^\tau \) are the components of the non-Abelian charge at site \( m \).\(^1\) The generators of left and right gauge transformations on the link \( L_{m,\ell}^\tau \) and \( R_{m,\ell}^\tau, \tau \in \{x, y, z\}, \) can be interpreted as color-electric fields, and the color-electric flux energy is \( J_m^\tau = \sum_\ell L_{m,\ell}^\tau R_{m,\ell}^\tau = \sum_\ell R_{m,\ell}^\tau R_{m,\ell}^\tau \). Now the Hilbert space of the \( m \)-th link is analogous to that of a quantum rigid rotator with total angular momentum \( J_m^\tau \), with body-fixed angular momentum operators \( L_{m,\ell}^\tau \) and space-fixed \( R_{m,\ell}^\tau \), and the basis elements can, thus, be labeled by the eigenvalues of \( J_m^\tau, L_{m,\ell}^\tau \) and \( R_{m,\ell}^\tau \), as \( |j_m^\ell e_m^{\rho} \rangle \). For the fermions, the Fock basis for each site now specifies the occupation numbers of each of the two color components, \( |n_m^1, n_m^2 \rangle \).

### 4. Numerical LGT calculations with TNS

The study of the models presented in section 3 with MPS techniques requires some previous considerations.

- The MPS formulation presented in Section 2.1 requires a finite dimensional Hilbert space for every site of the system. Although the gauge degrees of freedom do not fulfill this requirement, it is possible to truncate the basis of each link to a finite size. Numerical evidence suggests that at least for low energy states, a small truncation will be enough, even close to the continuum limit \([28]\).
- TNS can work with fermionic degrees of freedom without running into a sign problem, or increasing the computational complexity of the algorithms \([29, 30, 31]\). But the anticommutation of the corresponding operators requires specific attention in each step of the algorithm. In the one dimensional case, this can be simplified by applying a Jordan-Wigner transformation that maps the fermions to spins, which then can be treated by the most standard MPS algorithms. This strategy is adopted by the works we discuss next.

\(^1\)If present, external charges also need to be included.
• Only MPS that satisfy Gauss law are acceptable ansätze for physical states. The constraint can be imposed as a local symmetry at the level of the individual tensors [32, 33], such that only gauge invariant TNS are included in the ansatz. This can be generalized to higher dimensions [16, 34, 17, 35]. However, in the one dimensional case there is also the possibility to completely integrate out the gauge degrees of freedom, as they are completely fixed by the fermionic content, and to formulate the problem in the strictly physical space, thus dealing with the problem in a most efficient way in terms of number of variables, but at the cost of introducing non-local terms in the Hamiltonian. Notice that in this case, no truncation of the gauge degrees of freedom is needed, in principle [36, 37].

Over the last few years, different numerical studies have approached the problems above with various choices for boundary conditions and symmetries, and have systematically analyzed the continuum limits and the errors involved. As a consequence, the abundance of nowadays available results demonstrates beyond doubt the feasibility of the MPS ansatz for the physical states of gauge theories. In the following subsections, we review concrete numerical results that illustrate some of the most significant steps in this program.

4.1 Spectral properties

4.1.1 Mass gaps

![Figure 2: Dispersion relations for the massive Schwinger model. The left panel shows the results for open boundary MPS in the physical subspace [36], for fermion mass $m/g = 0.125$, system size $N = 160$ and lattice spacing given by $x = 1/(ga)^2 = 25$. The colors indicate the ground state (red), vector (green) and scalar (blue) branches, and symbols correspond to bond dimensions $D = 40$ (crosses), 80 (circles), and 100 (dots). The right plot [32] shows the vector excitation energies for non-zero momenta, computed using the uMPS ansatz and already converged in bond dimension, for $m/g = 0.75$, and lattice spacings $x = 100, 300, 800$ (circles), as well as the continuum estimates (stars). The curves fit the data to the relativistic dispersion relation $E^2 = k^2 + M^2(x)$. Right figure reproduced with permission from [32], @2014 APS.

The MPS ansatz can be used to approximate the lowest energy eigenstates of the lattice model, and after taking the appropriate continuum limit, calculate the particle content of the continuum theory. In the case of the Schwinger model, the results can be benchmarked against the analytical
solution at zero fermion mass, to assess the performance of the TNS calculations. The first impressive result was obtained early on in [11] using the original DMRG algorithm and, compared to previous numerical results, improved by several orders of magnitude the precision of the ground state energy and first (vector) mass gap in the continuum for massive and massless fermions. But only the TNS formulation allowed the power of DMRG to be extended for higher excitations, time evolution and other problems. More recently, in [36], using finite MPS with open boundary conditions in the physical space, and in [32], using a translationally and gauge invariant uMPS, calculations of the vector and scalar mass gaps were performed with unprecedented precision. In [32], the second vector excitation was accessed numerically for the first time.

Each MPS simulation is run for a finite lattice spacing and with a certain maximum bond dimension. In the case of finite systems as in [36], also for a fixed system size. For each such case, the algorithm identifies the ground state and several excitations, which differ in quantum numbers (e.g. left panel of figure 2). To reach the continuum limit, several extrapolation steps are necessary.

1. Bond dimension $D$: increasing the size of the tensors in the simulation improves the precision systematically. In the case of the Schwinger model, low energy states indeed converge very fast in $D$, as illustrated by the left plot in figure 2, where data for $D = 40, 80, 100$ collapse on top of each other for a system of $N = 160$ sites.\(^2\)

2. System size $N \to \infty$. This is necessary only in case of finite system simulations, which have in exchange the advantage of being unbiased, and able to cope with non-translationally-invariant scenarios. The uMPS works directly in the thermodynamic limit and does not require this step, but it assumes a local form for the excitations, and may find worse convergence if they are extended, as is the case for very small fermion mass.

3. Lattice spacing $x \to \infty$. To approach the continuum limit for a fixed fermion mass, simulations with large enough values of $x$ (and the corresponding adimensional masses $\mu$) are required. In the continuum limit, the theory becomes critical, and in principle harder for the MPS algorithms. Yet, the large values of $x$ (correspondingly small lattice spacings) reachable with MPS, as compared to Monte Carlo calculations, allow for more precise determination of the continuum values than with any other numerical technique, in the general case.\(^3\) It is actually possible to discriminate the presence of cubic or other higher order corrections in the lattice spacing.

The analysis shows that the main source of error is the continuum limit, a systematic uncertainty that can, nevertheless, be taken into account by a standard error analysis procedure, similar to those in data analysis for Lattice QCD [36]. The extremely precise estimations of the mass spectrum obtained with different MPS algorithms and for different fermion masses (far from the perturbative regime) are shown in table 1.

A similar analysis has also been performed for the 1+1 dimensional SU(2) Hamiltonian [37]. In that case, the continuum theory becomes massless as $m/g \to 0$, and the MPS simulations allow a study of critical exponents.

\(^2\)These numbers are to be compared to the largest possible bond dimension for a system of this size, $D_{\text{max}} = 2^{80}$.

\(^3\)For the massless fermion case, the most precise results have been obtained using strong coupling expansion [38].
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Table 1: Binding energies of the vector and scalar particles and their reported errors obtained with open boundary finite MPS with gauge degrees of freedom integrated out (left columns) or gauge invariant uniform MPS (right column) simulations. For the massless fermion, the results can be compared to the analytical values for the vector and scalar binding energies, respectively $M_V/g = 0.5641895$ and $M_S/g = 1.12838$.

<table>
<thead>
<tr>
<th>$m/g$</th>
<th>$M_V/g$</th>
<th>$M_S/g$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.56421(9)</td>
<td>0.56418(2)</td>
</tr>
<tr>
<td>0.125</td>
<td>0.53953(5)</td>
<td>0.539491(8)</td>
</tr>
<tr>
<td>0.25</td>
<td>0.51922(5)</td>
<td>0.51917(2)</td>
</tr>
<tr>
<td>0.5</td>
<td>0.48749(3)</td>
<td>0.487473(7)</td>
</tr>
</tbody>
</table>

4.1.2 Chiral condensate

From the MPS optimized for a certain state of the spectrum, e.g. the vacuum, any local or few body observable can be efficiently computed. This allows the study of different quantities in the continuum limit. One such study has been performed for the fermionic condensate, $\langle \bar{\Psi}(x)\Psi(x)\rangle/g$, the order parameter of chiral symmetry breaking. This quantity is UV divergent for massive fermions, the divergence coming solely from the free ($g = 0$) theory. The algorithms mentioned in the previous section [39, 40] also explored this observable in the massless and massive regimes. The precision of the results allowed for extracting the divergent behavior and identifying the leading corrections with the lattice spacing, and produced the most precise estimates of the chiral condensate in the ground state for non-vanishing fermion masses. Similar results have also been obtained with the related iDMRG algorithm [41].

4.1.3 The entropy of the vacuum

Figure 3: Entanglement entropy of the half chain in the vacuum of a lattice gauge theory. The leftmost panel shows the extrapolation to the continuum in the case of the Schwinger model, for $m/g = 0.25$ [42]. The four plots on the right correspond to the SU(2) model [37] and show, for $m/g = 0.8$, the different contributions to the entanglement entropy for each possible cut of the chain (a), as well as the bond dimension (b), system size (c) and continuum (d) extrapolations. Left figure reproduced with permission from [42], @2016 APS.

Of particular interest is the numerical analysis of the bipartite entanglement entropy of the
vacuum, a quantity that can be accessed very efficiently in MPS. On the one hand, in the continuum limit, which corresponds to a massive relativistic QFT, the entanglement entropy is predicted to diverge as \( S = (c/6) \log_2(\xi/a) \) \[43\], where \( c \) is the central charge of the conformal field theory describing the system at the critical point. The MPS data accurately capture this behavior, as has been explicitly checked for the Schwinger \[42\] and the SU(2) \[37\] models (see figure 3), where the numerical data were fitted to the logarithmic divergence, and the value of \( c \) was extracted. In \[42\], this allowed the definition of a renormalized entropy difference in the presence of external charges. On the other hand, the definition of the entanglement entropy of a gauge invariant lattice system entails some subtleties, since the gauge constraints are not local with respect to a straightforward bipartition of the Hilbert space \[44, 45\]. Instead, different contributions to the entropy can be identified, of which only one is distillable, and the others arise from the gauge invariance of the state. With the MPS ansatz, it is possible to compute these different contributions (see right part of figure 3) and to study their scaling and dependence on the parameters of the model.

4.2 Thermal properties

![Figure 4: Temperature dependence of the chiral condensate in the continuum for massless fermions, from finite size MPS on the physical space \[46\] (left) and gauge invariant uMPS (right, reproduced from Ref. \[47\]). Notice both studies used an opposite change convention. In both cases, the analytical results for the restoration of chiral symmetry (solid line) are recovered with good accuracy. On the left plot, two sets of data are compared, obtained using different approximations for the imaginary time evolution operator.

Using the MPO ansatz for the density operator it is possible to study temperature dependent properties of LGT. This has been applied to extract the thermal evolution of the chiral condensate in the massless and massive Schwinger model \[46, 47\]. Analytical calculations for \( m/g = 0 \) predict a smooth restoration of the chiral symmetry at high temperatures, a feature that the MPS calculations reproduce with good fidelity (figure 4).

The approximation of thermal equilibrium states is usually done by evolving the thermofield double from infinite temperature to the desired \( \beta = 1/T \). This introduces an additional parameter, namely the width of the imaginary time step, which has to be taken into account in the extrapolations towards the continuum. When using Suzuki-Trotter approximation this is nevertheless a well-controlled error and, as in the mass gap calculations, the largest uncertainty source is found to be the continuum extrapolation. This explains why the quality of the numerical results worsens towards infinite temperature: while the MPO approximation for the thermal state of a given spin
chain is actually easier at high temperature, much smaller lattice spacings are needed to correctly capture the lattice effects in that regime.

MPS allow the same calculations also for massive fermions and over the whole range of temperatures, reaching regimes for which no previous precise numerics existed [48, 47].

### 4.3 Finite density

![Figure 5: Phase diagram of the multiflavor Schwinger model, from [49]. The left plot shows the continuum extrapolation of the isospin number for the case of massless fermions, as a function of the (normalized) difference of chemical potentials and for different physical volumes. The steps show the location of the transitions, which can be compared to the analytical prediction (vertical lines). The right plot shows the phase diagram in the plane chemical potential difference vs. fermion mass, for fixed volume $L_g = 8$, with regions of different colors representing phases characterized by different isospin number.](image)

One of the most prominent features of TNS techniques is that they are free from the sign problem. The first explicit demonstration of this capability for a lattice calculation including the continuum limit extrapolation has been performed in [49] for the Schwinger model with two fermion flavors and non-vanishing chemical potential defined in Eq. (3.2).

The case of massless fermions can be solved analytically in the continuum, and at fixed volume exhibits a rich structure of phases, characterized by the *isospin number* $\Delta N$ (imbalance between fermion number for both flavors) and separated by first order phase transitions. These features are recovered from the MPS calculations carried out at constant finite physical volume, as shown in the left panel of figure 5. By performing the same analysis for massive fermions (right panel of the figure), the phase diagram can be explored beyond the Monte Carlo accessible regimes.

### 4.4 Real time evolution

Another interesting aspect of TNS methods is the possibility to simulate out of equilibrium dynamics. In the case of the massive Schwinger model, the first real time simulations have already been performed with MPS [32, 50]. Although these studies did not include a full continuum extrapolation as the ones above, the simulations were carried out close to the continuum (at very small lattice spacing, $x = 100$). The results allowed a study of the Schwinger pair production mechanism for quenches of varying strength, well beyond the linear response regime and, for moderate times, found effects that escape other approximate real-time methods for the lattice.
5. Discussion and perspectives

This paper has focused on studies where the TNS techniques are seen as a numerical tool, alternative to standard LGT methods, for solving continuum quantum field theories. The excellent results achieved for one dimensional problems have demonstrated beyond doubt the validity of the ansatz for the states of the theory, and its ability to capture all the relevant physics, also in scenarios that present difficulties for standard methods. A more ambitious goal is to extend these simulations to higher dimensional TNS, as PEPS. Such generalization is not trivial, but the recent algorithmic advances [51, 52], the development of gauge invariant formulations [16, 17, 34, 35], and the success of the one dimensional calculations constitute a solid motivation to pursue this endeavor.

As advanced in the introduction, there are other research directions connecting TNS and LGT. First of all, it is also possible to use a tensor network to represent the path integral for a lattice gauge model in space-time, and apply TNS algorithms to approximate the contraction without suffering from a sign problem, even for two spatial dimensions (see e.g. [53, 54, 55, 56]). Another active area of research is using TNS to study LGT that do not necessarily have a continuum limit, such as quantum link models [57]. Successful simulations have been performed for one-dimensional U(1) and SU(2) models for static and dynamical quantities [33, 58, 59]. A topic closely related to this is the possible quantum simulation of gauge theories in the lab using ultracold atoms [60]. Actually a first pioneering experiment has already been performed for the quantum simulation of the Schwinger model using trapped ions [61]. The TNS requirements mentioned above, namely a Hamiltonian formulation, and the finite-dimensional Hilbert space of local degrees of freedom, are also necessary for a quantum simulation with such systems, and TNS methods are optimally suited to assist in the design and verification of experimental protocols.

Finally, while comparably precise TNS simulations in higher dimensions can take some time to be realized, in the meantime we can expect that TNS help uncovering interesting features of other kinds of quantum field theories, even in one spatial dimension (see e.g. [62] in this volume), similar to the way in which DMRG has enabled great advances in condensed matter problems.

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